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	(71) Applicant: WHITEHRAD INSTITUTE FOR B CAL RESEARCH [US/US]; Nine Cambridg Cambridge, MA 02142 (US).	IOME ge Cen	DI- ter,		
٠	(72) Inventors: LIN, Herbert, Y.; 550 Memorial D 12D1, Cambridge, MA 02139 (US). WANG, X 872 Massachusetts Avenue, Ne 401, Cambri 02139 (US). WEINBERG, Robert, A.; 25 Cop Brookline, MA 02146 (US). LODISH, Harvey Fisher Avenue, Brookline, MA 02146 (US).	iao-Fa idge, A ley Str	Π V(A oct.		
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TGF-6 TYPE RECEPTOR CDNAs AND USES THEREFOR

Description

Background

Transforming growth factor-beta (TGF-8) is a member 5 of a family of structurally related cytokines that elicit a variety of responses, including growth, differentiation, and morphogenesis, in many different cell types. (Roberts, A.B. and M.B. Sporn, In: Peptide Growth Factors and Their Receptors, Springer-Verlag, Heidelberg, 10 pp. 421-472 (1990); Massague, J., Annu. Rev. Cell. Biol. 6:597-641 (1990)) In vertebrates at least five different forms of TGF- β , termed TGF- β 1 to TGF- β 5, have been identified; they all share a high degree (60%-80%) of amino-acid sequence identity. While TGF-\$1 was initially 15 characterized by its ability to induce anchorageindependent growth of normal rat kidney cells, its effects on most cell types are anti-mitogenic. (Altschul, S.F. et al., J. Mol. Biol. 215:403-410 (1990); Andres, J.L. et al., J. Cell. Biol. 109:3137-3145 (1989)). It is 20 strongly growth-inhibitory for many types of cells, -including both normal and transformed epithelial, endothelial, fibroblast, neuronal, lymphoid, and hemato-: poietic cells. In addition, TGF- β plays a central role in regulating the formation of extracellular matrix and 25 cell-matrix adhesion processes.

In spite of its widespread effects on cell phenotype and physiology, little is known about the biochemical mechanisms that enable $TGF-\beta$ family members to elicit these varied responses. Three distinct high-affinity

cell-surface TGP-β-binding proteins, termed type I, II and III, have been identified by incubating cells with radiolabelled TGF-β1, cross-linking bound TGP-β1 to cell surface molecules, and analyzing the labelled complexes by polyacrylamide gel electrophoresis. (Massague, J. and B. Like, J. Biol. Chem. 260:2636-2645 (1985); Cheifetz, s. et al. J. Biol. Chem. 261:9972-9978 (1986).) The binding constants are about 5-50pM for the type I and II receptor and 30-300 pM for the type III receptor. (Boyd, F.T. and J. Massague, J. Biol. Chem. 264:2272-2278

The type I and II receptors, of estimated 53 and 70-100 kilodaltons mass respectively, are N-glycosylated transmembrane proteins that are similar in many respects. Each of these receptors has a distinct affinity for each 15 member of the TGF- β family of ligands. (Boyd, F.T. and J. Massague, J. Biol. Chem. 264:2272-2278 (1989)) In contrast, the type III receptor shows comparable affinities for all TGF- β isotypes; the type III receptor is the most abundant cell-surface receptor for TGP-\$\beta\$ in many 20 cell lines (upwards of 200,000 per cell), and is an integral membrane proteoglycan. It is heavily modified by glycosaminoglycan (GAG) groups, and migrates heterogeneously upon gel electrophoresis as proteins of 280 to 330 kilodaltons. When deglycosylated with heparitinase 25 and chondrontinase, the protein core migrates as a 100-110 kilodalton protein. The TGF-β binding site resides in this protein core, as non-glycosylated forms of this receptor that are produced in cell mutants defective in GAG synthesis are capable of ligand binding 30 with affinities comparable to those of the natural receptor. (Cheifetz, S. and J. Massague, J. Biol. Chem., 264:12025-12028 (1989) A variant form of type III

receptor is secreted by some types of cells as a soluble molecule that apparently lacks a membrane anchor. This soluble species is found in low amounts in serum and in extracellular matrix.

The type III receptor, also called betaglycan, has a 5 biological function distinct from that of the type I and II receptors. Some mutant mink lung epithelial cell (MviLu) selected for loss of TGF-β responsiveness no longer express type I receptors; others, similarly selected, lose expression of both the type I and II 10 receptors. However, all these variants continue to express the type III receptor. (Boyd, F.T. and J. Massague, J. Biol. Chem. 264:2272-2278 (1989); Laiho, M. et al., J. Biol. Chem. 265:18518-18524 (1990)) This has led to the proposal that types I and II receptors are 15 signal-transducing molecules while the type III receptor, may subserve some other function, such as in concentrating ligand before presentation to the bona fide signal-transducing receptors. The secreted form of type III receptor, on the other hand, may act as a reservoir 20 or clearance system for bioactive TGF-β.

Additional information about each of these TGF-\$\beta\$ receptor types would enhance our understanding of their roles and make it possible, if desired, to alter their functions.

25 Summary of the Invention

The present invention relates to isolation, sequencing and characterization of DNA encoding the TGF-β type III receptor of mammalian origin and DNA encoding the TGF-β type II receptor of mammalian origin. It also relates to the encoded TGF-β type III and type II receptors, as well as to the soluble form of each; uses

of the receptor-encoding genes and of the receptors themselves; antibodies specific for TGF- β type III receptor and antibodies specific for TGF- β type II receptor. In particular, it relates to DNA encoding the TGF- β type III receptor of rat and human origin, DNA encoding the TGF- β type III receptor of human origin and homologues of each.

The TGF- β receptor-encoding DNA of the present invention can be used to identify equivalent TGF- β receptor type III and type II genes from other sources, using, for example, known hybridization-based methods or 10 the polymerase chain reaction. The type III receptor gene, the type II receptor gene or their respective encoded products can be used to alter the effects of TGF- β (e.g., by altering receptivity of cells to TGF- β or interfering with binding of TGF- β to its receptor), such 15 as its effects on cell proliferation or growth, cell adhesion and cell phenotype. For example, the TGF- β receptor type III gene, the TGF- β receptor type II gene, or a truncated gene which encodes less than the entire receptor (e.g., soluble TGF- β type III receptor, soluble 20 TGF-β type II receptor or the TGF-β type III or type II binding site) can be administered to an individual in whom TGF- β effects are to be altered. Alternatively, the TGP- β type III receptor, the TGF- β type II receptor, a soluble form thereof (i.e., a form lacking the membrane 25 anchor) or an active binding site of the TGF-β type III or the type II receptor can be administered to an individual to alter the effects of TGF- β .

Because of the many roles TGF-β has in the body, availability of the TGF-β receptors described herein makes it possible to further assess TGF-β function utilizing in vivo as well as in vitro methods and to alter (enhance or diminish) its effects.

Brief Description of the Drawings

Figure 1 is the DNA sequence (SEQ ID NO. 1) and the translated amino acid sequence (SEQ ID NO. 2) of type III TGP-#1 receptor cDNA clone R3-OFF (full insert size 6 kb), in which the open reading frame with flanking sequences of the clone are shown. The transmembrane domain is indicated by a single underline. Peptide sequences from purified type III receptor, mentioned in text, that correspond to the derived sequence, are in italics and underlined. Potential N-linked glycosylation sites are indicated by #, and extracellular cysteines by &. A consensus protein kinase C phosphorylation site is indicated by \$. The last non-vector encoded amino acid of Clone R3-OF (2.9 kb) is indicated by @. Consensus proteoglycan attachment site is indicated by +++. Other 15 potential glycosaminoglycan attachment sites are indicated by +. The upstream in-frame stop codon (-42 to -44) is indicated by a wavy line. Signal peptide cleavage site predicted by vonHeijne's algorithm (von Heijne, G., Nucl. Acid. Res. 14:4683-4690 (1986) is 20 indicated by an arrow.

Figure 2 is the nucleotide sequence of the fulllength type II TGF-β receptor cDNA clone 3FF isolated from a human HepG2 cell cDNA library (full insert size 5 kb) (SEQ ID NO. 3). The cDNA has an open reading frame 25 encoding a 572 amino acid residue protein.

Figure 3 is the amino acid sequence of the fulllength type II TGF- β receptor (SEQ ID NO. 4).

Detailed Description of the Invention

The subject invention is based on the isolation and sequencing of DNA of vertebrate, particularly mammalian, origin which encodes $TGF-\beta$ type III receptor and DNA of mammalian origin which encodes $TGF-\beta$ type II receptor,

expression of the encoded products and characterization of the expressed products. As described, a full-length cDNA which encodes TGF-β receptor type III has been isolated from a cDNA library constructed from a rat vascular smooth muscle cell line and a full-length cDNA which encodes TGF-β type II receptor has been isolated from a human cDNA library. The human homologue of the type III gene has also been cloned. A deposit of human TGF-β type III cDNA in the plasmid pBSK has been made under the terms of the Budapest Treaty at the American Type Culture Collection (10/21/91) under Accession Number 75127. All restrictions upon the availability of the deposited material will be irrevocably removed upon granting of a U.S. patent based on the subject

Isolation and Characterization of TGF-β Type III Receptor

As described herein, two separate strategies were pursued for the isolation of the TGF- β type III receptor In one approach, monoclonal antibodies were generated against the type III receptor protein and used to purify the receptor, which was then subjected to microsequencing. (See Example 1) Microsequencing of several peptides resulting from partial proteolysis of 25 the purified receptor produced four oligopeptide sequences, which were used to construct degenerate oligonucleotides. The degenerate oligonucleotides were used either as primers in a cloning strategy using the polymerase chain reaction (PCR) or as probes in screening 30 cDNA libraries. Although this strategy did not prove to be productive, the oligopeptide sequences were useful in verifying the identity of the receptor clones isolated by the second strategy.

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In the second approach to isolating TGF- β receptorencoding clones, an expression cloning strategy was used in COS cells; direct visualization of receptor positive cells was used to isolate receptor cDNAs. (See Example 2) In this approach, a cDNA library was constructed from A-10 cells, a rat vascular smooth muscle cell line which expresses all three TGF- β receptors (type I, II and III). COS cells transfected with cDNA components of this library in a vector carrying the cytomegalovirus (CMV) transcriptional promoter and the SV40 origin of repli-10 cation were screened to identify cells expressing substantially higher than normal levels of TGF-# receptor. One transfectant expressing such high levels of a TGF- β binding protein was identified and the original pool of expression constructs from which it was derived was split 15 into subpools, which were subjected to a second round of screening. Two further rounds of sib-selection resulted in isolation of one cDNA clone (R3-OF) with a 2.9 kb insert which induced high levels of TGF-\$-binding proteins in approximately 10% of cells into which it was introduced. The specificity of the TGF- β binding was validated by showing that addition of a 200-fold excess unlabeled competitor TGF-\$1 strongly reduced binding of 125 I-TGF- β to transfected cells.

The R3-OF cDNA encoded an open reading frame of 817 amino acid residues, but did not contain a stop codon. R3-OF was used as a probe to isolate a full-length cDNA from a rat 208F library. The resulting clone, R3-OFF, is 6kb in length and encodes a protein of 853 amino acids, which is colinear with clone R3-OF. The nucleotide sequence of R3-OFF is shown in Figure 1, along with the translated amino acid sequence.

Characterization of the receptor encoded by R3-OFF was carried out, as described in Example 3. Results . showed three distinct TGF-β binding protein species of TGF-β on the surface of mock-transfected COS cells, which is in accord with results reported by others. (Massague, J. et al., Ann. NY Acad. Sci. 593:59-72 (1990)). These included the two lower molecular weight type I and II receptors (65 and 85 kD) and the higher molecular weight type III proteoglycan, which migrates as a diffuse band of 280-330 kd. Enzymatic removal of the proteoglycan yielded a core protein of approximately 100 kd. Binding to all three receptor types is specific in that it was competed by 200-fold excess of unlabeled TGF-β1.

Transfecting the isolated cDNA caused a two-fold increase in expression of the type III receptor. When a cell lysate derived from COS cells transfected with clone R3-OFF was treated with deglycosylating enzymes, the heterogeneous 280-330 kd band was converted to a protein core which co-migrates with the type III protein core seen in parental A10 cells. Importantly, the recombinant protein core migrated differently from the endogenous COS cell type III protein core.

These observations were confirmed and extended using stably transfected cells expressing the type III cDNA.

L6 rat skeleton muscle myoblasts do not express any detectable type III mRNA and no endogeneous surface type III receptor (Massague et al., 1986; Segarini et al., 1989). These cells were transfected with the isolated cDNA in the vector pcDNA-neo. Cell clones stably expressing this clone in both the forward and reverse orientations with respect to the CMV promoter were isolated and analyzed by ligand binding assay.

Introduction of either the full-length clone R3-OFF or the partial clone R3-OF in the forward orientation resulted in expression of type III receptor. L6 cells transfected with the cDNA clones in the reverse orientation did not express this protein. Importantly, the apparent size of the protein core of the type III receptor in cells transformed with the R3-OF clone is smaller than that from R3-OFF transformed cells, consistent with the difference in the sizes of the protein cores predicted from their nucleic acid sequences.

Surprisingly, binding of radio-labeled ligand to the type II receptor was increased by 2.5 fold in cells expressing the type III cDNA. Binding to the type I receptor was unchanged. This apparently specific up-regulation of ligand-binding to the type II receptor was evident in all of the 15 stably transfected L6 cell lines analyzed to date. Furthermore, this effect seems to be mediated equally well by the full-length clone or a truncated clone (R3-OF) that lacks the cytoplasmic domain of TGF-\$\beta\$ type III receptor was expressed.

Expression of type III receptor mRNA was assessed by Northern blot analysis and RNA blot analysis. Northern gel analysis showed that the type III receptor mRNA is expressed as a single 6 kb message in several rat

25 tissues. RNA dot blot analysis of several different tissue culture cell lines was also carried out. Cells of mouse origin (MEL and YH16) appear to express a smaller (~5.5 kb) message for the type III mRNA than those of pig, rat and human origin. In all of these cells,

30 expression or absence of the type III mRNA is consistent with the expression or absence of detectable cell surface

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type III receptors, with the notable exception of the retinoblastoma cell lines (Y79, Weri-1, Weri-24, and Weri-27). These cells lack detectable surface expression of type III receptor, which confirms an earlier report. (Kimchi, A. et al., Science 240:196-198 (1988)). It is striking that the type III receptor mRNA is expressed in these cells at a level comparable to that of other cells that do indeed express type III receptor proteins at readily detectable levels. It appears that $TGF-\beta$ receptor III expression, which is substantial in normal retinoblasts (AD12), has been down-regulated in these retinoblastoma tumor cells, perhaps through post-transcriptional mechanisms.

The nucleotide sequence full reading frame along with flanking sequences of the full-length cDNA clone R3-OFF was determined and is presented in Figure 1. The reading frame encodes a protein of 853 amino acid residues, which is compatible with the 100 kD size observed for the fully deglycosylated TGF-\$1 type III receptor. The identity of the receptor as $TGF-\beta$ type III 20 was verified by searching for segments of the putative transcription product which included the peptide sequences determined by microsequencing of the isolated type III receptor. (See Example 1) As indicated in Figure 1, two segments of derived protein (underlined and 25 italicized, residues 378-388 and 427-434) precisely match with the amino acid sequences of two peptides (I and III) determined from direct biochemical analysis of the purified type III receptor.

Further analysis showed that TGF-\$ type III binding protein has an unusual structure for a cytokine receptor. Hydropathy analysis indicates that the protein includes a

N-terminal signal sequence, followed by a long, hydrophilic N-terminal region. A 27 residue region of strong hydrophobicity (underlined in Figure 1, residues 786-812) toward the C-terminus represents the single putative transmembrane domain. This suggests that nearly all of the receptor which is an N-terminal extracellular domain is anchored to the plasma membrane near its C-terminus. A relatively small C-terminal tail of 41 residues represents the cytoplasmic domain.

Analysis of related sequences provides few clues to 10 function of TGF-β type III protein. Only one other gene described to date, a glycoprotein expressed in high quantities by endothelial cells and termed endoglin, contains a related amino acid sequence. The most homologous regions between the sequences of the type III 15 receptor and endoglin (74%) falls primarily in the putative transmembrane and cytoplasmic domains. Similar to the general structure of type III receptor, endoglin is a glycoprotein which contains a large hydrophilic N-terminal domain which is presumably extracellular, 20 followed by a putative transmembrane domain and a short cytoplasmic tail of 47 amino acid residues. The biological role of endoglin is still unclear at present, although it has been suggested that it may involved in cell-cell recognition through interactions of an "RGD" 25 sequence on its ectodomain with other adhesion molecules. Unlike the TGF- β type III receptor, endoglin does not carry GAG groups.

Isolation of TGF- β Type II Receptor

The cDNA encoding the type II TGF-β receptor was also isolated, using expression cloning in COS cells. A full-length cDNA (designated clone 3FF) was isolated by high stringency hybridization from a human HepG2 cell cDNA library. (See Example 6) Analysis showed that the corresponding message is a 5 kb message which is expressed in different cell lines and tissues. Sequence analysis indicated that the cDNA has an open reading frame encoding a core 572 amino acid residue protein.

The nucleotide sequence of the full-length type II TGF-β receptor cDNA clone 3FF is shown in Figure 2; the amino acid sequence is represented in Figure 3.

The 572 amino acid residue protein has a single putative transmembrane domain, several consensus glycosylation sites, and a putative intracellular serine/ threonine kinase domain. The predicted size of the encoded protein core is -60 kd, which is too large for a type I TGF-β receptor. Instead, crosslinking experiments using iodinated TGF-β and COS cells transiently transfected with clone 3FF shows over-expression of a protein approximately 70-80 kd which corresponds to the size of type II TGF-β receptors. Thus, clone 3FF encodes a protein that specifically binds TGF-β and has an expressed protein size of 70-80 kd, both characteristic of the type II TGF-β receptor.

Uses of the Cloned TGF-β Receptors and Related Products

For the first time, as a result of the work described herein, DNAs encoding two of the three high affinity cell-surface TGF-β receptors have been isolated, their sequences and expression patterns determined and

the encoded proteins characterized. Expression of the $TGF-\beta$ type III receptor in cells which do not normally express the receptor, followed by ligand binding assay, verified that the cloned type III receptor-encoding DNA (i.e., either the full-length clone R3-OFF or the partial clone R3-OF) encoded the receptor. In addition, the work described herein resulted in the surprising finding that binding of $TGF-\beta$ to type II receptors in cells expressing the type III DNA was increased by 2.5 fold.

Additional insight into the role of the TGF- β type 10 III receptor and its interaction with TGF- β type II receptor is a result of the work described. For example, the role of TGF-8 type III receptor is unclear, but it has been proposed that it serves a most unusual function of attracting and concentrating TGF-\$s for eventual transfer to closely situated signal-transducing receptors. While most cytokines bind to a single cell surface receptor, members of the TGF- β family bind with greater or lesser affinity to three distinct cell surface proteins. This has raised the question of why these 20 three receptors are displayed by most cell types and whether they subserve distinct functions. Evidence obtained to date suggests that the type III receptor may perform functions quite different from those of types I and II. Thus, type III is substantially modified by GAGs 25 while types I and II appear to carry primarily the N-linked (and perhaps O-linked) sidechains that are characteristic of most growth factor receptors. addition, variant cells that have been selected for their ability to resist TGF- β -induced growth inhibition show 30 the absence of Type I or Type II receptors while continuing to display Type III receptors. Together, these data have caused some to propose that the Type I

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and II receptors represent <u>bona fide</u> signal-transducing receptors while the type III receptor, described here, plays another distinct role in the cell.

It remains possible that the type III receptor serves a most unusual function of attracting and concentrating TGF-\$\beta\$s on the cell surface for eventual transfer to closely situated signal-transducing receptors. Such a function would be unprecedented for a proteinaceous receptor, although heparin sulfate has been shown to activate basic FGF by binding to this growth factor prior to FGF association with its receptor (Yayon, A. et al., Cell 64:841-848 (1991)) Parenthetically, since the type III receptor also contains large quantities of heparan sulfate side-chains, it may also bind and present basic FGF to its receptor.

Evidence that is consistent with the role for the type III receptor comes from the work with L6 rat myoblast cells which is described herein. As described above, in L6 cells overexpressing type III receptor, the binding of radiolabelled TGF- β to the type II receptor is increased several fold when compared with that seen with parental cells. Further assessment of TGF- β type III function and interaction with type II and type I receptors will be needed to answer these questions and can be carried out using the materials and methods described here.

TGF- β receptors, both type III and type II, can be identified in other species, using all or a portion of the DNA encoding the receptor to be identified as a probe and methods described herein. For example, all or a portion of the DNA sequence encoding TGF- β type III receptor (shown in Figure 1) or all or a portion of the

DNA sequence encoding TGF-\$ type II receptor (shown in Figure 2) can be used to identify equivalent sequences in other animals. Stringency conditions used can be varied, as needed, to identify equivalent sequences in other species. Once a putative TGF-\$ receptor type III or type II-encoding sequence has been identified, whether it encodes the respective receptor type can be determined using known methods, such as described herein for verification that the cDNA insert of full-length clone R3-OFF and the cDNA insert of partial clone R3-OF encode 10 the type III receptor. For example, DNA isolated in this manner can be expressed in an appropriate host cell which does not express the receptor mRNA or the surface receptor (e.g., L6 rat skeleton muscle myoblasts) and analyzed by ligand binding (TGF-\$ binding) assay, as described herein. 15

Also as a result of the work described herein, antibodies (polyclonal or monoclonal) specific for the cloned TGF-\$ type III or the clones TGF-\$ type II receptor can be produced, using known methods. Such 20 antibodies and host cells (e.g., hybridoma cells) producing the antibodies are also the subject of the present invention. Antibodies specific for the cloned TGF-8 receptor can be used to identify host cells expressing isolated DNA thought to encode a TGF-8 25 receptor. In addition, antibodies can be used to block or inhibit TGF-\$\beta\$ activity. For example, antibodies specific for the cloned TGF-8 type III receptor can be used to block binding of TGF-8 to the receptor. They can be administered to an individual for whom reduction of TGP- β binding is desirable, such as in some fibrotic diseases (e.g., of skin, kidney and lung).

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The method of the present invention can be used for diagnosis of disorders involving abnormal binding of . TGF- β to TGF- β type III receptors and/or TGF- β type II receptors, such as fibrotic diseases. Abnormal binding of TGF- β to TGF- β type III receptor or TGF- β type II receptor at a cell surface may be measured, resulting in a test binding value, which is compared to an appropriate control binding value. Control binding values can be obtained using control cells known to have abnormal binding of TGF- β to its receptors or control cells which 10 are normal cells (e.g., evidence TGF- β binding to the TGP-6 receptor is within physiological levels). Control values are obtained by determining the extent to which TGF- β binds the appropriate receptor (i.e., TGF- β type III receptor or TGF-β type II receptor); such values can 15 be obtained at the time the test binding value is determined or can be previously determined (i.e., a previously determined standard). A test binding value similar to the control binding value obtained from abnormal cells is indicative of abnormal binding of TGF- β 20 to TGF-β type III receptor or TGF-β type II receptor. A test binding value similar to the control binding value obtained from normal cells is indicative of normal binding of TGF- β to TGF- β type III receptor or TGF- β type II receptor.

DNA and RNA encoding TGP- β type III receptor and DNA and RNA encoding TGF- β type II receptor are now available. As used herein, the term DNA or RNA encoding the respective TGF-\$ receptor includes any oligodeoxynucleotide or oligodeoxyribonucleotide sequence 30 which, upon expression, results in production of a TGF- β receptor having the functional characteristics of the

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TGF- β receptor. That is, the present invention includes DNA and RNA which, upon expression in an appropriate host cell, produces a TGP- β type III receptor which has an affinity for TGF- β similar to that of the TGF- β type III receptor on naturally occurring cell surfaces (e.g., it shows comparable affinities for all TGP- β isotypes). Similarly, the present invention includes DNA and RNA which, upon expression in an appropriate host cell, produces a TGF- β type II receptor which has an affinity for TGF- β similar to that of TGF- β type II receptor on 10 naturally occurring cell surfaces (e.g., it has a distinctive affinity for each member of the TGP- β family of ligands similar to that of the naturally occurring TGF- β type II receptor). The DNA or RNA can be produced -in an appropriate host cell or can be produced 15 synthetically (e.g., by an amplification technique such as PCR) or chemically.

The present invention also includes the isolated TGF-β type III receptor encoded by the nucleotide sequence of full-length R3-OFF, the isolated TGF-β type III receptor encoded by the nucleotide sequence of partial clone R3-OF, the isolated TGF-β type II receptor encoded by the nucleotide sequence of full-length clone 3FF and TGF-β type III and type II receptors which bind TGF-β isotypes with substantially the same affinity. The isolated TGF-β type III and type II receptors can be produced by recombinant techniques, as described herein, or can be isolated from sources in which they occur naturally or synthesized chemically. As used herein, the terms cloned TGF-β type III and cloned TGF-β type II

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described herein, and TGF- β type III and type II receptors (e.g., from other species) which exhibit . substantially the same affinity for the TGF-\$\beta\$ isotypes as the respective receptors.

As described previously, cells in which the cloned TGF- β type III receptor is expressed bind TGF- β in essentially the same manner as do cells on which the type III receptor occurs naturally. Further analysis of ligand interactions with the cloned TGF- $oldsymbol{eta}$ type III receptor, based upon site-directed mutagenesis of both $TGF-\beta$ and the receptor, can be carried out to identify residues important for binding. For example, DNA having the sequence of Figure 1 can be altered by adding, deleting or substituting at least one nucleotide, in order to produce a modified DNA sequence which encodes a modified cloned TGF- β type III receptor. The functional characteristics of the modified receptor (e.g., its TGF- β -binding ability and association of the binding with effects normally resulting from binding) can be assessed, using the methods described herein. Modification of the cloned TGF-β type III receptor can be carried out to 20 produce, for example, a form of the TGF-β type III receptor, referred to herein as soluble TGF-# receptor, which is not membrane bound and retains the ability to bind the TGF- β isotypes with an affinity substantially 25 the same as the naturally-occurring receptor. Such a TGF-8 type III receptor could be produced, using known genetic engineering or synthetic techniques; it could include none of the transmembrane region present in the naturally-occurring TGF- β type III receptor or only a small portion of that region (i.e., small enough not to

interfere with its soluble nature). For example, it can include amino acids 1 through 785 of the TGF- β type III sequence of Figure 1 or a portion of that sequence sufficient to retain TGF- β binding ability (e.g., amino acids 24-785, which does not include the signal peptide cleavage site present in the first 23 amino acids). A soluble TGF- β type II receptor (e.g., one which does not include the transmembrane and cytoplasmic domains) can also be produced. For example, it can include amino acids 1 through 166, inclusive, of Figure 3 or a sufficient portion thereof to retain TGF- β binding ability substantially the same as that of TGF- β type II receptor.

The TGP-β type III receptor and/or type II receptor can be used for therapeutic purposes. As described above, 15 the TGF-β family of proteins mediate a wide variety of cellular activities, including regulation of cell growth, regulation of cell differentiation and control of cell metabolism. TGF- β may be essential to cell function and most cells synthesize TGF- β and have TGF- β cell surface 20 receptors. Depending on cell type and environment, the effects of TGF-\$\beta\$ vary: proliferation can be stimulated or inhibited, differentiation can be induced or interrupted and cell functions can be stimulated or suppressed. TGF- β is present from embryonic stages 25 through adult life and, thus, can affect these key processes throughout life. The similarities of a particular TGF-β (e.g., TGF-β1) across species and from cell to cell are considerable. For example, the amino acid sequence of a particular $TGF-\beta$ and the nucleotide sequence of the gene which encodes it regardless of

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source, are essentially identical across species. This further suggests that $TGF-\beta$ has a critical role in essential processes.

Specifically, TGF-\$\beta\$ has been shown to have antiinflammatory and immune suppression capabilities, to play an important role in bone formation (by increasing osteoblast activity), inhibit cancer cell proliferation in culture, and control proliferation of glandular cells of the prostate. As a result, it has potential therapeutic applications in altering certain immune system 10 responses (and possibly in modifying immune-mediated diseases); in treating systemic bone disease (e.g., osteoporosis) and conditions in which bone growth is to be enhanced (e.g., repair of broken bones) and in controlling growth and metastasis of cancer cells. In 15 addition, TGF- β appears to play a role in determining whether some cell types undergo or do not undergo mitosis. In this respect, TGF- β may play an important role in tissue repair. Some diseases or conditions appear to involve low production or chronic overproduction of TGF- β . (For example, results of animal studies suggest that there is a correlation between the over production of TGF-\$\beta\$ and diseases characterized by fibrosis in the lung, kidney, liver or in viral mediated immune expression.)

Clearly, TGF-\$\beta\$ has key roles in body processes and numerous related potential clinical or therapeutic applications in wound healing, cancer, immune therapy and bone therapy. Availability of TGF-\$\beta\$ receptor genes, the encoded products and methods of using them in vitro and in vivo provides an additional ability to control or

regulate TGF- β activity and effect in the body. Por example, the TGP-\$ type II or type III receptor encoded by the type II or the type III receptor genes of the subject invention can be used, as appropriate, to alter the effects of TGF-# (e.g., to enhance the effect of TGF-\$\beta\$ in the body or to inhibit or reduce (totally or partially) its effects). It is also possible to administer to an individual in whom TGF-\$ bound to TGF-\$ type III receptor, such as soluble TGF- β type III receptor. The present invention provides both a TGF-\$ agonist and a 10 TGF-# antagonist. For these purposes, DNA gene encoding the entire TGF-\$ type II or type III receptor, the encoded type II or type III receptor or a soluble form of either receptor can be used. Alternatively, antibodies or other ligands designed based upon these sequences or specific for them can be used for this purpose.

Knowledge of the amino acid sequences of TGF-β type
III and type II receptors makes it possible to better understand their structure and to design compounds which interfere with binding of the receptor with TGF-β. It
20 makes possible identification of existing compounds and design of new compounds which are type III and/or type II receptor antagonists.

Cells expressing the type III and/or type II receptors of the present invention can be used to screen compounds for their ability to interfere with (block totally or partially) TGF binding to the receptors. For example, cells which do not express TGF-β type III receptor (e.g., L6 rat skeleton muscle myoblasts) but have been modified to do so by incorporation of the type III cDNA in an appropriate vector can be used for this

purpose. A compound to be assessed is added, for example, to tissue culture dishes containing type III. expressing cells, along with labeled TGP- β . As a control, the same concentration of labeled TGF- β is added to tissue culture dishes containing the same type of cells. After sufficient time for binding of TGF- β to the receptor to occur, binding of labeled TGF-\$ to the cells is assessed, using known methods (e.g., by means of a gamma counter) and the extent to which it occurred in the 10 presence and in the absence of the compound to be assessed is determined. Comparison of the two values show whether the test compound blocked TGF- β binding to the receptor (i.e., less binding in the presence of the compound than in its absence is evidence that the test 15 compound has blocked binding of TGF- β to the TGF- β type III receptor).

Alternatively, a cell line expressing the TGF-β receptor or cells expressing microinjected TGF-β receptor RNA, can be used to assess compounds for their ability to block TGF-β binding to the receptor. In this embodiment, a compound to be assessed is added to tissue culture dishes containing the cell line cells expressing microinjected TGF-β receptor RNA, along with TGF-β. As a control, TGF-β alone is added to the same type of cells expressing microinjected endothelin receptor RNA. After sufficient time for binding of TGF-β to the receptor to occur, the extent to which binding occurred is measured, both in the presence and in the absence of the compound to be assessed. Comparison of the two values shows whether the compound blocked TGF-β binding to the receptor. The TGF-β type III and type II receptors can

also be used to identify $TGF-\beta$ -like substances, to purify $TGF-\beta$ and to identify $TGF-\beta$ regions which are responsible for binding to the respective receptors. For example, the type III receptor can be used in an affinity-based method to identify substances which bind the receptor in a manner similar to $TGF-\beta$.

The invention will now be illustrated by the following examples, which are not intended to be limiting in any way.

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EXAMPLES

Materials and methods used in Examples 1-5 are described below.

Materials

The following is a description of materials used in the work described herein.

Recombinant human TGF-\$1 was provided by Rik Derynck of Genentech. COS-M6 cells were provided by Brian Seed of the Massachusetts General Hospital and Alejandro Aruffo of Bristol-Myers-Squibb. Heparitinase was provided by Tetsuhito Kojima and Robert Rosenberg of MIT.

LLC-PK1 cells were a gift of Dennis Ausiello of the Massachusetts General Hospital. YH-16 cell were a gift of Edward Yeh of the Massachusetts General Hospital. 3-4 cells were a gift of Eugene Kaji of the Whitehead

25 Institute for Biomedical Research. All other cell lines were purchased from ATCC and grown as specified by the supplier, except as noted.

Expression Cloning

Construction of cDNA Library and Generation of ...
Plasmid Pools

10µg polyadenylated mRNA was prepared from A10 cells by the proteinase-K/SDS method (Gonda et al., Molec. Cell. Biol. 2:617-624 (1982)). Double stranded cDNA was synthesized and linkered to nonpalindromic BstX1 adaptors as described by Seed, B. and A. Aruffo, Proc. Natl. Acad. Sci. USA 84:3365-3369 (1987). Acaptored cDNA was sizefractionated on a 5 to 20% potassium acetate gradient, and inserts greater than 1 kb were ligated to the plasmid vector pcDNA-1, and electroporated in the \underline{E} . \underline{coli} MC1061/P3, yielding a primary library with a titer of >107 recombinants. A portion of the cDNA was plated as 15 pools of -1x104 recombinant bacteria colonies grown on 15 cm petri dishes with Luria-broth agar containing 7.5 mg/ml tetracycline and 12.5 mg/ml ampicillin. Cells were scraped off the plates in 10 mls of Luria-broth, and glycerol stocks of pooled bacteria were stored at -70°C. The remaining bacteria was used to purify plasmid DNA using the alkaline lysis mini-prep method (Sambrook, J. et al., Molecular Cloning: A Laboratory Manual, 2d Ed. (Cold Spring Harbor, NY, Cold Spring Harbor Laboratory Press (1989)).

Plasmid pools (each representing -1x10⁴ clones) were transfected into COS-M6 (subclone of COS-7 cells) by the DEAE-dextran/chloroquine method described by Seed, B. and A. Aruffo, Proc. Natl. Acad. Sci. USA 84:3365-3369

(1987). Forty-eight hours after transfection, cells were

incubated with 50 pM123I-TGF-\$1 (100 to 200 Ci/mmol) for
4 hours at 4°C), autoradiographic analysis of transfected
cells was performed using NT-B2 photographic emulsion
(Kodak) essentially as described (Gearing, D.P. et al.,
EMBO J. 8:3667-3676 (1989)). After development of
slides, cells were air-dried and mounted with mounting
media and a glass coverslip. Slides were analyzed under
an Olympus OM-T1 inverted phase-contrast microscope using
a dissection trans-illuminator for darkfield illumination.

Subdivision of Positive Pool

of 86 pools screened, one pool (#13) was identified as positive and a glycerol stock of bacteria corresponding to this pool was titered and 25 pools of 1000 clones were generated and miniprep plasmid from these pools were transfected into COS cells as described above. Several positive pools of 1000 were identified, and one was replated as 25 plates of 100 colonies. A replica was made of this positive plate and harvested. Once a positive pool was identified, individual colonies were picked from the corresponding master plate and grown overnight in 3 ml liquid culture. A 2-dimensional grid representing the 100 clones was generated and a single clone, R3-OF, was isolated.

25 Cloning of R3-OFF

A 208F rat fibroblast library in lambda ZAP II (Stratagene) was screened at high stringency with clone R3-OF insert, and several clones with -6kb inserts were isolated, one of which is referred to as R3-OFF.

DNA Sequencing and Sequence Analysis

Double-stranded DNA was sequenced by the dideoxy. chain termination method using Sequenase reagents (United States Biochemicals). Comparison of the sequence to the data bases was performed using BLAST (Altschoul, S.F. et al., J. Mol. Biol. 215:403-410 (1990)).

Iodination of TGF-β

TGF-\$1 was iodinated using the chloramine-T method as described (Cheifetz, S. and J.L. Andres, <u>J. Biol.</u>
Chem. 263:16984-16991 (1988)).

Chemical Cross-Linking

Transfected COS cells grown on 10 cm dishes or subconfluent L6 and A-10 cells grown on 3.5 cm dishes were incubated with 125I-TGF-β1 in binding buffer 15 (Frebs-Ringer buffered with 20 mM Hepes, pH 7.5, 5 mM MgSO4, 0.5% BSA), washed 4 times with ice-cold binding buffer without BSA, and incubated for 15 minutes with binding buffer without BSA containing 60ng/ml disuccinimidyl suberate at 4°C under constant rotation. 20 Crosslinking was terminated by addition of 7% sucrose in binding buffer. Cells were scraped, collected and pelleted by centrifugation, then resuspended in lysis buffer (10 mM Tris, pH 7.4, 1 mM EDTA, pH 8.0, 1% Triton-X 100, 10 μ g/ml of pepstatin, 10 μ g/ml leupeptin, 25 10 μ g/ml antipain, 100 μ g/m; benzamidine hydrochloride, 100 μ g/ml soybean trypsin inhibitor, 50 μ g/ml aprotonin, and 1 mM phenylmethylsulfonyl fluoride). Solubilized material was analyzed by 7% SDS-PAGE and subjected to

autoradiographic analysis by exposure to X-AR film (Kodak) at -70°C.

Enzymatic Digestion

Digestion of solubilized TGF-b receptors with chondroitinase and heparitinase was performed as described (Cheifetz, S. and J.L. Andres, J. Biol. Chem. 263:16984-16991 (1988); Segarini, P.R. and S.M. Seyedin, J. Biol. Chem., 263: 8366-8370 (1988).

Generation of Stable Cell Lines

L6 myoblasts were split 1:10 into 10 cm dishes and 10 . transfected the following day by the calcium phosphate method (Chen, C. and H. Okayama, Molec. Cell. Biol. 7:2745-2752 (1987)) with clones R3-OF or R3-OFF in the forward and reverse orientations in the vector pcDNA-neo 15 (Invitrogen). Cells were subjected to selection in the presence of G418 (Geneticin, GIBCO) for several weeks until individual colonies were visible by the naked eye. These clones were isolated and amplified.

RNA Blot Analyses

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Rat tissue polyadenylated mRNA was prepared by the lithium chloride/urea method (Auffrey, C. and F. Raugeon, Eur. J. Biochemistry 107:303-313 (1980), followed by oligo-dT cellulose chromatography (Aviv and Leder, 1972). Polyadenylated mRNA from cell lines was prepared by the 25 proteinase K/SDS method (Gonda, T.J. et al., Molec. Cell. Biol. 2:617-624 (1982)). Samples of mRNA were resolved by electrophoresis on 1% agarose-2.2M formaldehyde gels, blotted onto nylon membranes (Biotrns, ICN) and incubated with the 2.9 kb insert of clone Re-OF labeled with ³²P by random priming as probe (Sambrook, J. et al., Molecular Cloning: A Laboratory Manual, 2d Ed., Cold Spring Harbor, NY, Cold Spring Harbor Laboratory Press, (1989). Hybridizations were performed at 42°C in hybridization buffer containing 50% formamide overnight, and blots were washed at 55°C in 0.2X SSC, 0.1% SDS, before exposure to X-AR film at -70°C.

Example 1. Production of Anti-Type III Receptor Protein

Antibodies and Microsequencing and Microsequencing of Peptides Resulting from Partial

Proteolysis of Purified Type III Receptor

Initially cellular proteoglycans were purified from human placenta and then subjected to enzymatic deglycosy-15 lation with heparitinase and chondroitinase. Protein cores in the molecular weight range of 100-130 kilodaltons were further purified by preparative gel electrophoresis; these should include the type III receptor. This partially purified material was used as an immunogen 20 in mice. After screening 850 hybridoma lines developed from immunized mice, three lines were found to produce antibodies that specifically recognized and immunoprecipitated a deglycosylated polypeptide species of 100-120 kD. This species could be radiolabelled by incubation of whole cells with 125 I-TGF-β followed by covalent cross-linking. Its size is consistent with that of the protein core previously reported for the type III receptor. (Massague, J., Annu. Rev. Cell. Biol. 6:597-641 (1990))

Monoclonal antibody 94 was used to purify the type III receptor from rat liver by affinity-chromatography. The purified receptor was subjected to partial proteolysis and the resulting peptides were resolved by high pressure liquid chromatography. Several peptides were subjected to microsequencing and yielded the following oligopeptide sequences:

Peptide I: ILLDPDHPPAL (SEQ ID NO. 5)

Peptide II: QAPFPINFMIA (SEQ ID NO. 6)

10 Peptide III: QPIVPSVQ (SEQ ID NO. 7)

Peptide IV: FYVEQGYGR (SEQ ID NO. 8)

These peptide sequences were used to construct degenerate oligonucleotides that served either as primers in a cloning strategy using the polymerase chain reaction (PCR) or as probes in screening cDNA libraries. While this strategy was not productive, the oligopeptide sequences proved useful in verifying the receptor clones isolated by the second, alternative strategy (see Example 2).

20 Example 2. Expression Cloning of the Type III Receptor cDNA

An expression cloning strategy in COS cells, a procedure which takes advantage of the considerable amplification of individual cDNAs in transfected COS cells was used as an alternative means to isolate TGF-β receptor clones. Such amplification is mediated by SV40 large T antigen expressed by the COS cells interacting

with a SV40 origin of replication in the cDNA vector.

Gearing, D. et al., EMBO J. 8:3667-3676 (1989); Lin,

H.Y., et al., Proc. Natl. Acad. Sci. 88:3185-3189

(1991a); Lin, H. Y. et al., Science, in press (1991);

Mathews, L. S. and Vale, W. W., Cell 65:973-982 (1991).

The strategy involved the construction of a cDNA library from A-10 cells, a rat vascular smooth muscle cell line that expresses all three high-affinity TGF- β The resulting cDNAs were inserted into the receptors. vector pcDNA-1, which carries the CMV transcriptional 10 promoter and the SV40 origin of replication. The resulting library was then divided into pools of 10,000 independent recombinants each and DNA from each pool was transfected into 1.5 x 10 COS-7 cells grown on glass flaskettes by means of DEAE-dextran transfection proce-15 dure. Aruffo, A. and Seed, B., Proc. Natl. Acad. Sci., U.S.A. 84:8573-8577 (1987); Gearing, D. et al., EMBO J. 8:3667-3676 (1989); Mathews, L. S. and Vale, W. W., Cell 65:973-982 (1991). The transfected cells were cultured for 48-60 hours and then exposed to radiolabelled TGF- β 1 20 for four hours. Following this treatment, the glass slides carrying these cells were washed extensively and fixed. These slides were dipped in liquid photographic emulsion and examined by darkfield microscopy. While all of the receptor genes cloned to date by this procedure 25 have undetectable or low constitutive levels of expression in COS cells, we were hindered by the fact that untransfected COS cells already express substantial amounts of type III TGF-\$ receptor. Such expression, estimated to be approximately 2 x 105 receptor molecules per cell, might well have generated an unacceptably high level of background binding. However, since the detection procedure involves visualizing radiolabelled

ligand-binding on individual cells, it was hoped that identifying occasional cells expressing substantially higher levels of vector-encoded receptor would be possible. This hope was vindicated in the initial experiments.

After screening nearly one million cDNA clones in this manner, a glass slide containing 20 positive transfectants was identified. The original pool of expression constructs from which one such transfectant was derived was split into 25 subpools of 1000 clones each and these were subjected to a second round of screening. Two further rounds of sib-selection resulted in the isolation of a cDNA clone (R3-0F) with a 2.9 kb insert that induced high levels of TGF-β-binding proteins in approximately 10% of COS cells into which it was transfected.

The specificity of this binding was validated by showing that addition of a 200-fold excess of unlabeled TGF-β competitor strongly reduced binding of 125 I-TGF-β to transfected cells. By taking into account a transfection efficiency of 10% and the high background of endogenous receptor expression, we calculated that the level of total 125 I-TGF-β binding to each glass slide of cells transfected with this cDNA clone (Figure 1C) was only 2-fold above the level seen with mock transfectants (data not shown). Nonetheless, this marginal increase in ligand-binding was sufficient to identify rare transfectants amidst a large field of cells expressing this background level of receptor.

The R3-OF cDNA encoded an open reading frame of 836 amino acid residues of which the 3' most 18 were encoded by vector sequence, clearly indicating that clone R3-OF

was an incomplete cDNA insert which ended prematurely at the codon specifying alanine 818 (Figure 4). R3-OF was used as a probe to isolate a full-length cDNA from a rat 208F lambda phage library. This clone, termed R3-OFF, was 6 kb in length and encoded a protein of 853 amino acids; its sequence was co-linear with that of clone R3-OF.

Example 3. Characterization of the Product of the Full Length Clone R3-OFF

Characterization of the product of the full length clone R3-OFF was undertaken in order to determine which of the three TGF-β receptors it specified. To do so, COS transfectants were incubated with radioiodinated TGF-β, chemical crosslinker was added and the labelled receptors were resolved by polyacrylamide gel electrophoresis.

Labelling of cell surface TGF-β receptors in this way resulted in the detection of three distinct species on the surface of COS cells, as extensively by others (Massague, J. et al., Ann. NY Acad. Sci. 593:59-72 (1990). These included the two lower molecular weight type I and II receptors (65 and 85 kD) and the higher molecular weight type III proteoglycan, which migrated as a diffuse band of 280-330 kd. Enzymatic treatment of the proteoglycan with chondroitinase and heparitinase yielded a core protein of approximately 100 kd. Binding to all three receptor types was specific, in that it was completed by 200-fold excess of unlabeled TGF-β1.

Transfecting the R3-OFF cDNA caused a two-fold increase in expression of the type III receptor. When a cell lysate derived from COS cells transfected with clone

R3-OFF was treated with deglycosylating enzymes, the heterogenous 280-330 kd band was converted to a protein core which co-migrated with the type III protein core seen in untransfected A10 cells. Importantly, the recombinant protein core migrates differently from the endogenous COS cell type III protein core.

These observations were confirmed and extended in experiments using stably transfected cells expressing the R3-OFF cDNA. L6 rat skeleton muscle myoblasts normally do not express detectable type III mRNA or endogenous type III receptor (determined by radiolabelled ligand-binding assay). Such cells were transfected with the isolated cDNA in the vector pcDNA-neo. Cell clones stably expressing this clone in both the forward and reverse orientations with respect to the CMV promoter were isolated and analyzed by ligand-binding assay.

Introduction of either the full length clone R3-OFF or the partial clone R3-OF in the forward orientation led to the de novo expression of the type III receptor. L6 cells transfected with the cDNA in reversed orientation did not express this protein. The apparent size of the protein core of the type III receptor in cells transfected with the R3-OF clone is smaller than that expressed by R3-OFF transfected cells, consistent with the difference in the sizes of the protein cores predicted from the respective nucleic acid sequences (Figure 1).

Unexpectedly, the amount of radio-labelled ligand corss-linked to the type II receptor is increased by 2.5 fold in cells expressing the type III cDNA, while the amount cross-linked to the type I receptor remained

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unchanged. This apparent specific up-regulation of ligand-binding to the type II receptor could be detected with all of the 15 stably transfected L6 cell lines analyzed so far. This effect seems to be mediated by the truncated clone R3-OF which lacks the cytoplasmic domain as well as by the full-length clone R3-OFF.

Example 4. Expression of Type III Receptor

Northern blot analysis demonstrated that the type
III receptor mRNA is expressed as a single 6 kb message
in several rat tissues. The level of mRNA expression in
the liver was less than in other tissues, a result
expected from earlier surveys of various tissues using
radioiodinated TGF-β1. Based on this information, it
appears that clone R3-OFF, with a -6 kb cDNA insert,
represents a full length rat type III cDNA clone.

cells of mouse origin (MEL and YH16) appear to express a smaller (~5.5 kb) message for the type III mRNA than those of pig, rat and human origin. In all of these cells, expression or absence of the type III mRNA is consistent with the expression or absence of detectable cell surface type III receptors with the notable exception of the retinoblastoma cell lines (Y79, Weri-1, Weri-24, and Weri-27). These cells have previously been shown to lack detectable surface expression of type III receptor, a result confirmed by our own unpublished work. It is striking that the type III receptor mRNA is expressed in these cells at a level comparable to that of other cells that do indeed express type III receptor proteins at readily detectable levels. At this moment, we can only conclude that TGF-β receptor III expression,

which is substantial in normal retinoblasts (AD12), has been down-regulated in these retinoblastoma tumor cells, perhaps through post-transcriptional mechanisms.

Example 5. Sequence Analysis of the Full-Length Type III cDNA

The full-length cDNA clone (R3-OFF), described in Example 4, was subjected to sequence analysis. The full reading frame along with flanking sequences is presented in Figure 1. This reading frame encodes a protein of 853 amino acid residues, which is compatible with the 100 kD observed for the fully deglycosylated TGF-β type III receptor.

Two segments of derived protein sequence (underlined and italicized, residues 378-388 and 427-434) precisely
15 match those determined earlier from direct biochemical analysis of the purified receptor protein. This further secured the identity of this isolated cDNA clone as encoding the rat type III receptor.

This TGF-\$\beta\$ binding protein has an unusual structure

for a cytokine receptor. Hydropathy analysis indicates a

N-terminal signal sequence, followed by a long,
hydrophilic N-terminal region (Kyte, J. and R. F.

Doolittle, J. Mol. Biol. 157:105-132 (1982)). A 27
residue region of strong hydrophobicity (underlined,
residues 786-812) toward the C-terminus represents the
single putative transmembrane domain. This suggests that
nearly all of the receptor is composed of an N-terminal
extracellular domain that is anchored to the plasma
membrane near its C-terminus. A relatively short

C-terminal tail of 41 residues represents the putative cytoplasmic domain.

The clone R3-OF was also analyzed and found to be a truncated version of R3-OFF, with an identical open reading frame but whose last encoded residue is alanine 818 (Figure 1).

In R3-OFF there are six consensus N-linked glycosylation sites and 15 cysteines (indicated in Figure 1). There is at least one consensus glycosaminoglycan addition site at serine 535 (Bernfield, M. and K. C. Hooper, Ann. N.Y. Acad. Sci. in press (1991), and numerous Ser-Gly residues that are potential sites for GAG conjugation. A consensus protein kinase C site is also present at residue 817.

Only one other gene described to date, a 15 glycoprotein expressed in high quantities by endothelial cells and termed endoglin (Gougos and Letarte, 1990), contains a related amino acid sequence. Overall, there is -30% identity with the type III receptor over the entire 645 amino acid residue length of endoglin. The 20 most homologous regions between the sequences of the type III receptor and endoglin (74% identity) falls primarily in the putative transmembrane and cytoplasmic domains. Similar to the general structure of type III receptor, endoglin is a glycoprotein which contains a large 25 hydrophilic and presumably extracellular N-terminal domain followed by a putative transmembrane domain and a short cytoplasmic tail of 47 amino acid residues. The biological role of endoglin is unclear, though it has been suggested that it may involve cell-cell recognition 30 through interactions of an "RGD" sequence on its

ectodomain with other adhesion molecules. Unlike the TGF-8 type III receptor, endoglin does not carry GAG groups.

Equivalents

Those skilled in the art will recognize, or be able to ascertain using not more than routine experimentation, many equivalents to the specific embodiments of the invention described herein. Such equivalents are intended to be encompassed by the following claims.

International Application No: PCT/

MICROOR	GANISMS
Outland Sheet in connection with the microorganism relayed to an	page 6 has 12 of the description i
A. IDENTIFICATION OF DEPOSIT	
Further deposits are identified on an additional chaet 0	<u> </u>
Name of depository institution 4	,
American Type Culture Colle	
Address of depository institution (including postal code and country)	•
12301 Parklawn Drive Rockville, Maryland 20852	USA
Date of deposit 6	Assuration Number 6
21 October 1991	75127
B. ADDITIONAL INDICATIONS 1 Coors blank if not applicable	which a European Patent is sought.
grant of the European Patent or un	withdrawn or is deemed to be biological material deposited with on under Accession No. e of a sample to an expert nominated be European Rule 28(5).
D. SEPARATE FURNISHING OF INDICATIONS 6 Gave bles The indicatests hand below will be automated to the international "Accession Sumber of Depart.")	uk if not espicable) I Burees later 9 (Specify the general nature of the indications s.s.,
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-m	(Authorized Officer)

Form PCT/RO/134 (January 1981)

CLAIMS

- Isolated DNA encoding TGF-β receptor of vertebrate origin or DNA which hybridizes thereto and encodes TGF-β receptor of vertebrate origin.
- 2. Isolated DNA of Claim 1 wherein the TGF- β receptor is TGF- β type III receptor or TGF- β type II receptor.
 - 3. Isolated DNA of Claim 2 which is of mammalian origin.
- 10 4. Isolated DNA of murine or human origin encoding $TGF-\beta$ type III receptor or DNA which hybridizes thereto.
- Isolated DNA of Claim 4 having the nucleotide sequence of Figure 1 or a portion thereof sufficient to encode TGF-β type III receptor.
 - 6. Isolated DNA of murine or human origin encoding TGF-β type II receptor or DNA which hybridizes thereto.
- Isolated DNA of Claim 6 having the nucleotide
 sequence of Figure 2 or a portion thereof sufficient to encode TGF-β type II receptor.
 - 8. Isolated TGF- β type III receptor of mammalian origin.

- 9. Isolated TGP- β type III receptor of Claim 8 having the amino acid sequence of Figure 1 or a substantially similar amino acid sequence.
- 10. Isolated TGF- β type II receptor of mammalian origin.
- 5 11. Isolated TGF- β type II receptor of Claim 10 having the amino acid sequence of Figure 3 or a substantially similar amino acid sequence.
 - 12. Recombinant TGF- β type III receptor of mammalian origin.
- 10 13. Recombinant TGF-β type III receptor of Claim 8 having the amino acid sequence of Figure 1 or a substantially similar amino acid sequence.
 - 14. Recombinant TGP-β type II receptor of mammalian origin.
- 15 15. Recombinant TGF- β type II receptor of Claim 10 having the amino acid sequence of Figure 4 or a substantially similar amino acid sequence.
 - 16. Soluble TGF-β receptor.
- 17. Soluble TGF-β receptor of Claim 16 which is soluble
 20 TGF-β type III receptor.
 - 18. Soluble TGF- β type III receptor of Claim 17 in which the amino acid sequence is amino acids 1 through

785, inclusive, of Figure 1 or a substantially similar amino acid sequence.

- 19. Soluble TGF- β receptor of Claim 16 which is soluble TGF- β type II receptor.
- 5 20. Soluble TGF-β receptor of Claim 19 in which the amino acid sequence is approximately amino acids 1 through 166, inclusive, of Pigure 3, or a substantially similar amino acid sequence.
- 21. An antibody which specifically recognized TGP-β type
 10 III receptor of mammalian origin.
- 22. An antibody of Claim 21 which is a monoclonal mantibody.
 - 23. An antibody which specifically recognizes soluble TGF-β type III receptor of mammalian origin.
- 15 24. An antibody which specifically recognizes soluble TGF-β type II receptor of mammalian origin.
- 25. A method of altering TGF-β binding to TGF-β type II or type III receptor on the surface of a cell, comprising combining soluble TGF-β type II or type III receptor with the cell, under conditions appropriate for binding of the soluble TGF-β receptor and TGF-β.

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- 26. The method of Claim 25 wherein TGF-β binding is inhibited.
- 27. A method of altering TGF-β binding to TGF-β type III receptor on the surface of a cell comprising combining the cell with DNA encoding TGF-β type III receptor in an appropriate expression system which expresses TGF-β type III receptor, under conditions appropriate for expression of TGF-β type III receptor and binding of TGF-β with TGF-β type III receptor.
- 28. A method of regulating the effect of TGF-β in a mammal, comprising administering to the mammal a TGF-β receptor selected from the group consisting of: TGF-β type III receptor, TGF-β type III receptor, soluble TGF-β type III receptor, soluble TGF-β type III receptor, TGF-β bound to TGF-β type III receptor or a combination thereof, in sufficient quantity to alter binding of TGF-β to TGF-β type III receptor, binding of TGF-β to TGF-β type III receptor or both.

- 29. TGF- β receptor according to any one of Claims 8 to 20, for use in therapy.
- 30. An antibody according to any one of Claims 21 to 24, for use in therapy.
- 31. Use of TGF- β receptor according to any one of Claims 8 to 20, for the manufacture of a medicament for altering (e.g. inhibiting) TGF- β binding to TGF- β type II or type III receptor on the surface of a cell.
- 10 32. Use of a TGF-β receptor selected from the group consisting of: TGF-β type III receptor, TGF-β type II receptor, soluble TGF-β type III receptor, soluble TGF-β type III receptor, TGF-β bound to TGF-β type III receptor or a combination thereof, for the manufacturing of a medicament for use in regulating the affect of TGF-β in a mammal.
 - 33. A method of assessing the ability of a compound to interfere with TGF- β binding to the TGF- β type III receptor, comprising the steps of:
- 20 a) combining:
 - 1) mammalian cells which express the $TGF-\beta$ type III receptor;
 - labeled TGF-β; and
 - 3) a compound to be assessed;

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- b) maintaining the product of (a) under conditions sufficient for TGF-β to bind to the TGF-β type III receptor;
- c) determining the extent of binding of TGF- β to TGF- β type III receptors in the presence of the compound to be assessed; and
- d) comparing the determination made in (c) with the extent to which binding of TGF-β to the TGF-β type III receptor occurs in the absence of the compound to be assessed,

wherein if TGF- β binding to the TGF- β type III receptor occurs to a lesser extent in the presence of the compound to be assessed than in the absence of the compound to be assessed, the compound to be assessed interferes with TGF- β binding to TGF- β type III receptors.

- 34. A method of Claim 33 wherein the cells which express the TGF- β type III receptor are a cell line.
- 35. A method of Claim 34 wherein the cells which express
 the TGF-β type III receptor are cells modified to
 express the TGF-β type III receptor.
 - 36. A method of Claim 35 wherein the cells modified to express the TGF- β type III receptor are cells which have incorporated into them TGF- β receptor cDNA in an appropriate vector or microinjected TGF- β receptor RNA.

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- 37. A method of assessing the ability of a compound to interfere with TGF- β binding to the TGF- β type II receptor comprising the steps of:
 - a) combining:
 - mammalian cells which express the TGF-β
 type II receptor;
 - labeled TGF-β; and
 - 3) a compound to be assessed;
 - b) maintaining the product of (a) under conditions sufficient for TGF- β to bind to the TGF- β type II receptor;
 - c) determining the extent of binding of TGF- β to TGF- β type II receptors in the presence of the compound to be assessed; and
 - d) comparing the determination made in (c) with the extent to which binding of TGF-β to the TGF-β type II receptor occurs in the absence of the compound to be assessed,
- wherein if TGF-β binding to the TGF-β type II

 receptor occurs to a lesser extent in the presence
 of the compound to be assessed than in the absence
 of the compound to be assessed, the compound to be
 assessed has interfered with TGF-β binding to TGF-β
 type II receptor.
- 25 38. A method of Claim 37 wherein the cells which express the TGF- β type II receptor are a cell line.
 - 39. A method of Claim 38 wherein the cells which express the TGF- β type II receptor are cells modified to express the TGF- β type II receptor.

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- 40. A method of Claim 39 wherein the cells modified to express the TGF-β type II receptor are cells which have incorporated into them TGF-β receptor cDNA in an appropriate vector or microinjected TGF-β receptor RNA.
- 41. A method of detecting abnormal binding of TGF- β TGF- β type III receptors of TGF- β type III receptors at a cell surface, comprising:
 - a) determining the extent of binding of TGF-β to TGF-β type III receptors or TGF-β type II receptors by cells in a sample obtained from an individual in whom binding is to be assessed thereby producing a test binding value; and
 - b) comparing the results of (a) with the extent to which binding occurs at the cell surface in control cells which are cells known to have abnormal binding of TGF-β to TGF-β type III receptors or TGF-β type II receptors resulting in a control binding value,
- wherein a test binding value similar to the control binding value is indicative of abnormal binding of $TGF-\beta$ to $TGF-\beta$ type III receptor or $TGF-\beta$ type II receptor.

FIGURE 1A

240													-	mc10	سين .	-180
AGG	AGGI	GA J	LAGTC	CCCC	coc	CTCC	GGAT	GGC	GCAG	FTG	CACT	GCCC	16 6		CTCGC	-120
				~~~	0 00	ርእሮጥ	നന	TCC	CCTA	GTA	ACTC	CICC	AC C			-110
			*	~2~	CTC	<u> </u>	CC NG	CCA	ACTT	GAA	CAGT	GCAG.	<b>~~</b> ^		TOTIV	- 00
AGC	TACA	cc (	<b>SACT</b>	TGCC	A CG	ATTG	CCTT	CYY	TCTG	AAG	AACC	AAAG	GC 1	GIIG	GAGAG	• •
										-		•				
																48
TG	GCA	GTG	ACA	TCC	CAC	CAC	ATG	ATC	ccc	GTG	ATG	GTT	GTC	CIG	N1G	16
iet .	Ala	Val	Thr	Ser	His	Hi=	Het	Ile	Pro	Val	Ket	Val	VAI	Leu	HOT	10
																96
GC	GCC	TGC	CTG	CCC	ACC	CCC	GGT	CCY	GAG	ccc	AGC	ACC.	2	101	Clu	32
Ser .	Ala	Сув	Leu	Ala	Thr	Ala	Gly	Pro	GIA	PIO	ser	Int	AFG	Cys	GIU	
•						Ť								•		
												mm^	3 mv	CAG	NCC.	144
CTG	LCY	CCY	ATC	AAC	GCC	TCT	CAC	CCA	GIC	CAG	33.5	116	NT.	Clu	Sor	48
Leu	Ser	Pro	Ile	Asn	λla	Ser	His	Pro	Val	GIN	VTG	rea	ne c	O10	361	•••
	ĵ.	•		#												
									101	ccc	200	300	ccc	CTG	CCA	192
rtc .	YCC	GTT	CTG Leu	TCT	GGC	TGT	GCC	AGC	AUA Aua	61.	Th-	The	Glv	Leu	Pro	64
Phe	Thr	Val	Leu	Ser	GIÅ	CyB	VTE	"Ser	vrA	GLY	1111	1.111	-LJ			
	`.			+		•										
			CAT	CTC	CT2	330	CTC	CGA	AGT	λCλ	GAT	CAG	GGA	CCA	GGC	240
AGG	GAG	GIC	His	610	Tan	lan	TAU	Ara	Ser	Thr	Ago	Gln	Glv	Pro	Gly	80
Arg	GIU	AST	HIB		Leu	Vell	Dec	y					4		•	
	~~~	010	AGA	CAG	CTT	ACC.	CTG	CAC	CTG	AAC	ccc	λTT	GCC	TCG	GTG	288
CAG	7	CAG	yrd	Gla	Val	Thr	Leu	His	Leu	Asn	Pro	Ile	Ala	Ser	Val	96
GTH	ALY		niy	GIU	***											
	3 Off	cin c	CAC	222	CCIP	ATC	GTG	TTC	CTG	CTC	AAC	TCC	ccc	CAG	ccc	336
CAC	ACT	UAU	His	Tura	Dro	Tle	Val	Phe	Leu	Leu	λsn	Ser	Pro	Gln	Pro	112
~~~	CTC	TCC	CAT	CTG	DAG	ACG	GAG	AGA	CTG	GCC	GCT	GGT	GTC	ccc	AGA	384
CIG	CIO	Too	His	TAU	Lvs	Thr	Glu	Arg	Leu	Ala	Ala	Gly	Val	Pro	λrg	128
CTC	***	CTC	CTT	TCG	GAG	GGT	TCT	GTC	GTC	CAG	TIT	CCA	TCA	GGA	AAC	432
701	Pho	Ter	Val	Ser	Glu	Glv	Ser	Val	Val	Gln	Phe	Pro	Ser	Gly	Asn	144
red	FIIG			-		,							+	•	#	
•																
mm/C	TOO	ምጥና	2 202	GCA	GAA	ACA	GAG	GAA	AGG	AAI	TTC	CCT	CAA	GAA	AAT	480
Dho	200	110	, Thr	Ala	Glu	Thr	Glu	Glu	Arq	Авг	Phe	Pro	Glr	Glu	Asn	160
Pne	Ser	Det		7.24					_							
CAF	Cha		T CHC	. ccc	TGG	GCC	CAA	AAG	GAA	TAT	C GGA	GCA	GTO	ACT	TCG	528
CIT	ut-	101	, Val	Arn	Tro	Ala	Gln	Lys	Glu	Tyi	Gly	Ala	Va]	l Thr	Ser	176
GIU	nle									•	•	-			•	
mmc		י מים	, (~T-C		ב ארם	GCA	AGP	AAC	: ATC	TAT	TTA 1	. AAA	GTO	GG?	CAA	576
TIC	ML.	. ci	n Yeu	, and		Ala	Arc	Asr	Ile	Ty	c Ile	Lys	Va.	l Gly	y Glů	192
rne	1111		ושנו	. Dyt	, ,,,,			,				•			-	

## PIGURE 1B

							•										<b>654</b>
	GAT	CAA	GTG	TTT	CCT	CCT	λCG	TGT	AAC	ATA	CCC	AAG	AAT	TTC	CTC	TCA	624
	Asp	Gln	Val	Phe	Pro	Pro	Thr	Cys	Asn	Ile	Gly	Lys	λsn	Phe	Leu	Ser	208
	•							E								٧,	•
	CTC	TAA	TAC	CTT	CCC	GAG	TAC	CTT	CYY	ccc	AAA	GCC	GCC	GAA	GGT	TGT	672 224
	Lou	Asn	Tyr	Leu	Ala	Glu	Tyr	Leu	Gln	Pro	Lys	Ala	YIE	GIA	GTÅ	Cyn E	224
																-	
												~~~	200	A TO	CAG	TTA	720
	GTC	CIG	ccc	AGT	CAG	ccc	CAT	GAA	AAG	GAA	GTA	CAC	ATC	TIA	Glu	Len	240
	Val	Leu	Pro	Ser	Gln	Pro	His	Glu	rys	GIA	ARI	810	114	110	O.L.O.	. 202	
								TAC	100	~~	T-100	CAG	GTG	CAT	ATA	ATA	768
	ATT	YCC	CCC	AGC	TCG	AAC	CCI	Tyr	SOT	21.	Phe	Gln	Val	ABD	Ile	Ile	256
	116	Thr.	PEO	Ser	ser	ABII	PIO	.131	361	~~		~					
	GTT	C > C	A TER	CC.	CCT	CCT	CAA	GAG	GAT	ccc	GAG	GTG	GTC	λλλ	AAC	CTT	816
	Ual	GWC	TIO	Ara	Pro	Ala	Gln	Glu	ABD	Pro	Glu	Val	Val	Lys	λsn	Leu	272
	Val	veh	110	9										-			
	GTC	CTG	ATC	TTG	AAG	TGC	AAA	λλG	TCT	GTC	AAC	TGG	GTG	ATC	AAG	TCT	864
	Val	Leu	Ile	Leu	Lys	Cys	Lys	Lys	Ser	Val	λsn	Trp	Val	Ile	Lys	Ser	288
					•	Ē	_										
													-				
an Sagrada si	TIT	GAC	GTC	AAG	GGA	YYC	TTG	УУУ	GTC	ATT	CCT	ccc	AAC	AGT	ATC	GGC	. 912
	Phe	Asp	Val	Lys	Gly	Asn	Leu	Lys	Val	Ile	Ala	Pro	Asn	Ser	116	GIY	. 304
																	960
	TTT	GGA	ÄAA	GAG	AGT	GAA	CGA	TCC	ATG	ACA	ATG	ACC	AAA	116	GIV	Ara	320
	Phe	Gly	Lys	Glu	Ser	Glu	Arg	Ser	Met	Thr	Met	Int	LYB	Leu	V=1	ar y	720
							~~~	GAG	220	OTIC	NTV:	336	TCG	GCA	CTG	GAC	1008
	GAT	GAC	ATC	CCT	TUU	MP	CAA	GAU	VVI	LAU	Met	T.VA	Tro	λla	Leu	Asp	336
	Asp	ABD	116	PIO	Set	1111	GIII	GLU	ADII	<i></i>		-,-					
	220	CCC		) ACC	CCA	CTC	ACG	TCA	TAC	ACA	ATG	GCT	ccc	GTG	GCT	λλτ	1056
	y av	Glv	Tur	Ara Ara	Pro	Val	Thr	Ser	Tvr	Thr	Het	λla	Pro	Val	Ala	Asn.	352
	ABU	413	-1-	**** 4	•••				-4-								
	AGA	TTT	CAT	CTT	CGG	CTT	GAG	AAC	AAC	GAG	GAG	ATG	AGA	GA1	: CAG	GAA	1104
	. Ara	Phe	His	Leu	Arq	Leu	Glu	Asn	Asn	Glu	Glu	Met	Arg	Asp	Glu	. Glu	368
٠.	GTC	CAC	ACC	: ATT	CCI	CCI	GAG	CTT	CGI	'ATC	CIG	CTG	GAC	: cc	GAC	CAC	1152
	Val	His	Thr	Ile	Pro	Pro	Glu	Leu	Arg	Ile	Lev	Lev	ABE	Pro	yet	His	384
			•								F	epti	.de 1				•
			·														
	CCG	ccc	: GCC	CIG	GAC	: AAC	CC	CIC	TTC	: cci	A GGA	GAC	GGZ	AG(	CC	TAA A	1200
	Pro	Pro	Ala	Lev	i yel	) Asr	Pro	Leu	Phe	Pro	o Gly	Gli	ı Çl	Se	Pro	nBA c	400
	,														~ i~~	2 BBC	1248
·.	GGT	GGI	CTC	ccc	TI	CC	TTO	cco	GA!	TAT	c cc	AGC	AGI	1 66	· Tu	G AAG	416
	Gly	Gly	Let	Pro	Phe	Pro	Phe	Pro	Asj	p 110	e pro	Arg	, Ar	J GI	A TE	p Lys	410
								. ~~-			0 034	2 CC	ጉ አጥረ	ი <u>ი</u> ლი	T CC	ር እርጥ	1296
	GAG	GGC	GA)	A GAT	AGO	AT(		. CGC		n AA	אניטי ט רח ה	, D-	S TI	o Ve	] Dr	C AGT o Ser	432
	Glu	Gl	Gli	ı Ası	o Ar	3 110	s Pr	o WEG	y rr	o ră	- 7T	S FI	ptid	- <del>Va</del>		o Ser	
												pe	PCIU	- 4			

# PIGURE 1C

									_:						ccc	1344
GTT	CAA	CTG	CTT	CCT	GAC	CAC	CGA	GAA	CCA	GAA	GAA	616	CAA	C)	27	448
Val	Gln	Leu	Leu	Pro	yab	His	Хrg	Glu	Pro	GIU	GIA	ATT	GIN	GIY	Gly <	. 440
GTG	GAC	ATC	GCC	CTG	TCA	GTC	AAA	TGT	GAC	CAT	Gλλ	AAG	ATG	GTC	GTG	1392
Val	Asp	Ile	Ala	Leu	Ser	Val	Lys	Cys	yab	His	Glu	Lys	Xet	Val	Val	464
	_							£						:		
GCT	GTA	GAC	λλλ	GAC	TCT	TTC	CAG	ACC	AAT	GGC	TAC	TCA	GGG .	ATG	GAG	1440
Ala	Val	GRA	Lvs	ABD	Ser	Phe	Gln	Thr	Asn	Gly	Tyr	Ser	Gly	Xet	Glu	480
				•								+				
СТС	ACC	CTG	TTG	GAT	CCT	TCG	TGT	λλλ	GCC	λλλ	ATG	λλΤ	GGT	λCT	CAC	1488
TOU	The	Len	T.OU	Ago	Pro	Ser	Cvs	Lys	Ala	Lys	Het	λsn	Gly	Thr	His	496
Ded	2111	200	200				3			•		#	•			
•							_									
mmm	077	OTC.	CNG	ጥርሞ	ccc	CTG	AAT	GGC	TGT	CCT	ACT	CGA	CAT	CCG	AGG	1536
111	011	Ton	Clu	202	Dro	1.011	Agn	Gly	Cva	Glv	Thr	Ara	His	λrq	λra	512
Pne	ATT	Dea	GIU	SEL	FLO	200	non	023	3	,		,				
									_							
		~~~	03 M	COR	CTC	OTT	ጥኔሮ	TAT	220	TCT	ATT	GTG	GŤG	CAG	GCT .	1584
TCU	ACC	700	CUT	601	Vai	Ual	Tur	Tyr	Agn	Ser	Tle	Val	Val	Gln	Ala	528
ser	Thr	PIO	мвр	CTA	ATT	AWI	ııı	-1-	NO.1	-						,
			~~~	~>m	300	N.C.	ccc	TCC	CCT	CAT	GCC	ሞልጥ	GAA	GAC	TTG -	1632
CCG	TCC	CCT	666	GAT	AGC.	wat	000	Trp	Dro	len	Gly	Tur	Glu	Aap	Leu	544
Pro	Ser	Pro	era	Авр	ser	261	GIĀ	Trp	FLO	veh	OLY	-1-	014	nop		
	• • •	-				***										
						-	~~i	GGA	C) C	ccc	CAT	GNA	CGA	GAA	ACT	1680
GAG	TCA	GGC	GAT	AAT	GUA	TII	CCI	COA	300	C1	Lan.	Clu	Gly	Glu	Thr	560
Glu	Ser	Gly	yab	Asn	GIĀ	rne	Pro	Gly	vab	GIA	vaħ	GIU	arl	910	****	
	+															
													100		ccc	1728
. GCC	ccc	CIG	AGC	CGA	GCT	GGA	GTG	GTG	GTG	TIT	AAC	160	AGC.	110	200	576
Ala	Pro	Leu	Ser	Arg	Ala	Gly	Val	Val	Val	Phe	Asn			rea	AFY	370
•											#	£				
																1776
CAG	CTG	AGG	AAT	ccc	AGT	GGC	TTC	CAG	GGC	CAG	CTC	GAT	GGA	AAT	GCT	
Gln	Leu	Arg	Asn	Pro	Ser	Gly	Phe	Gln	Gly	Gln	Leu	Asp	GIA	YBD	Yls	592
													•	#		
•																
ACC	TTC	AAC	: ATG	GAG	CTG	TAT	AAC	ACA	GAC	CTC	TTI	CIG	GTG	ccc	TCC	1824
Thr	Phe	Asn	Het	: Glu	Leu	Tyr	Asn	Thr	Asp	Leu	Phe	Leu	Val	Pro	Ser	608
•		•														
CCA	GGG	GTC	TTC	: TCI	GTG	GCA	GAG	AAC	GAG	CAT	GTI	TAI	GT1	GAG	GTG	1872
Pro	Glv	Val	Phe	e Ser	. Val	Ala	Glu	Asn	Glu	His	Va]	Tyr	· Val	Glu	Val	624
	_												•			
TCT	GTC	: ACC	AAC	GC1	GAC	: CA	GAT	CIG	GGA	TTC	: GCC	: ATC	CA)	ACC	TGC	1920
Sav	· Val	The	Lvi	Alz	Aso	Glr	Ast	Leu	Gly	Phe	Ala	ılle	Gli	The	Cys	640
J61	. · · · · ·	4	,												€.	

# PIGURE 1D

Lala	CTC	TCT	CCA	TAC	TCC	AAC	CCA	GAC	λGλ	ATG	TCT	GAT	TAC	YCC	ATC	1968
Phe	Leu	Ser	Pro	Tyr	Ser	Asn	Pro	λsp	λrg	Met	Ser	yab	Tyr	Thr	Ile	656
																2016
ATC	GAG	AAC	atc	TGT	CCG	УУУ	GAC	GAC	TCT	GTG	AAG	TTC	TAC	AGC	100	672
Ile	Glu	Asn	Ile	Cys	Pro	Lys	yab	ysb	Ser	AST	rys	rae	TYE	SAL		0,2
				£												
			~~~		000	3.000	~~	CAT	CCT	GAG	GTG	GAC	AAG	AAG	CGC	2064
AAG	AGA	UTG	UAC DIA	Dho	Pro	Tle	Pro	His	Ala	Glu	Val	λep	Lys	Lys	λrg	688
-																
TTC	AGC	TTC	CTG	TTC	AAG	TCT	GTG	TTC	AAC	ACC	TCC	CIG	CTC	TTC	CTG	2112
Phe	Ser	Phe	Leu	Phe	Lys	Ser	Val	Phe	Asn	Thr	Ser	Leu	Leu	Phe	Leu	704
									#							
											000	500		110	CENCE	2160
CAC	TGC	GAG	TTG	ACT	CTG	TGC	TCC	λGG	AAG	AAG	GUC	Ser	Leu	T.VE	Leu	720
Hib		Glu	Leu	Thr	Leu	Cys	ser	λrg	Lyu	Lyw	GLY	561	200	2,0		
	£					-										
ccc)CC	тст	GTG	ACT	CCT	GAC	GAC	GCC	TGC	ACT	TCT	CTC	GAT	GCC	ACC	2208
Pro	λra	CVB	Val	Thr	Pro	Asp	Asp	Ala	Cys	Thr	Ser	Leu	увр	Ala	Thr	736
	3					•			£							
٠																2256
ATG	ATC	TGG	ACC	ATG	ATG	CAG	AAT	AAG	AAG	ycy	TIC	ycc	AAG	CCC	CTG	752
Met	Ile	Trp	Thr	Met	Met	Gln	Asn	Lys	Lys	THE	PAG	TILL	Ly=	PLO	Leu	,,,
		oma.	CT(C	CNG	CTA	GAC	TAT	AAA	GAA	AAT	GTI	ccc	: AGC	ACT	λλG	2304
GCT	U10	Ual	Teu	Gln	Val	Asp	Tvr	Lvs	Glu	Asn	Val	Pro	Ser	Thr	Lys	768
														•		
GAT	TCC	AGT	CCA	ATT	CCI	CCT	CCT	CCT	CCA	CAG	ATT	TIC	CA1	. CCC	CTG	2352
Авр	Ser	Ser	Pro	Ile	Pro	Pro	Pro	Pro	Pro	Gln	Ile	Phe	e Hie	Gly	Leu	784
_	-		•													2400
GAC	λCG	CTC	ACC	GTG	λTG	GGC	ATT	GCA	TTI	GCX	GCX	TI	GIC	AIC	GGA	800
УвЪ	Thr	Leu	Thr	Val	Met	Gly	Ile	Ala	Phe	Ala	VIS	PRE	3 va.	116	Gly	000
			300			- T-T-C	TCC	TAC	እጥር	TAC	: TC	: CAC	e ac	A GGG	GAG	2448
GCG	CTC	CIG	The	· Glu	Als	Ten	TY	TVY	Tle	TVI	Sei	Hi	Th	c Gly	, Glu	816
ACA	GCA	. CGA	AGG	CAG	CAA	GTC	: cc1	: ACC	TC	cc	CCI	L GC	CTO	GAC	AAC	2496
Thr	Ala	Arc	Arc	Glr	Glr	val	Pro	Thr	Sex	Pro	Pro	Al:	a Se	r Gli	y yeu	832
\$		-		•												
																2544
AGC	AGC	: GCG	GCC	CAC	AGO	TA :	GGG	: AGC	AC:	r CA	G AG	r AC	_ D=	TG	C TCT	2544 · 848
Ser	Ser	Ala	Ala	His	s Sei	: Ile	e G1;	Ser	Th	r Gli	n se:	e In	r PI	o cy	s Ser	040
						~~m~	****	ChCI	, C) C	פרר ו	רניניי	CACC	GC A	GCCA	GGGCA	2599
						ou Tui	AUNE	GACI	TONC	المان			*1			853
ser	261	: Şei	Thi	. WI	2											

FIGURE 1E

,	DECCCCCATG	CCAGTGCTGC	GTGTCCACAG	TCAGAAGTCT	TGATCTGGGC	TCCCTGTAAA	2659
`,	AND COCOCA	ATTTCAGTAT	ACAGACAGCC	AGTTCTACCC	ACCCCTTACC	ACCCCCACA	2719
		CCTCCCCATC	TGTCACACGA	AAGCTAAGCT	GGTGGCCTTC	CCCACCAGCC	2779
	******	TOCCOCTTTC	AATGTGAAAC	ATCTGCCAGT	TITGTTTTGT	TTTTTTAATG	2839
	~~~	CACCTGTCCA	AACATCCATC	ATTTGGGGTG	GICTGITTIA	CAGAGTAAAG	2899
ì	CIGCILITIE	AGGGACGTCA	CCTACTGTGT	AGAGCCAAGG	GGAGACAGCT	AGGATTCTCG	2959
			AAATAGAAGA				2997

# FIGURE 2

				GTTGGCGAGG
		GCGCTGAGTT	GAAGTTGAGT	GAGTCACTCG
AGTTTCCTGT	TTCCCCCGCA	CCCCGCGCGT	GCACCCGCTC	GGGACAGGAG
CCCCCACCGA	GCGACGACAC	CCTCGGCCGC	CGGGGGCCTC	CCCGCGCCTC
CCGGACTCCT	GTGCAGCTTC	TGGCTGGCGA	GCGGGCGCCA	CATCTGGCCC
CCCCCCTCC	AGGCCCCTCC	CGCCGCGGG	TCCGGAGAGG	CCCCCCCCCC
GCACATCTGC	GCTGCCGGCC	GAAGGCGCCG	TCCGTGCGCT	GGGGGCTCGG
GAGCGCAGCC	AGGGGTCCGG	CTGCCATGGG	TCGGGGGCTG	CTCAGGGGCC
TCTATGACGA	GCAGCGGGGT	CTGTGGACGC	GTATCGCCAG	CACGATCCCA
TGTGGCCGCT	GCACATCGTC		ATGATAGTCA	CTGACAACAA
CCGCACGTTC	AGAAGTCGGT	TANTANCONC	ATTTTGTGAT	GTGAGATTTT
CCCTCCACTC	AAGTTTCCAC	TCCTGCATGA	GCAACTGCAG	CATCACCTCC
CCACCTGTGA	CAACCAGAAA	AGTOTOTOTO	GCTGTATGGA	GAAAGAATGA
ATCTGTGAGA	AGCCACAGGA	CAGTTTGCCA	TGACCCCAAG	CTCCCCTACC
CGAGAACATA	ACACTAGAGA	GCTGCTTCTC	CAAAGTGCAT	TATGAAGGAA
ATGACTITAT	TCTGGAAGAT	TTTCTTCATG	TGTTCCTGTA	GCTCTGATGA
AAAAAAAAGC	CTGGTGAGAC	TCTCAGAAGA	ATATAACACC	AGCANTCCTG
GTGCAATGAC	AACATCATCT	CAAGTGACAG	GCATCAGCCT	CCTGCCACCA
ACTTGTTGCT		CATCATCATC	TTCTACTGCT	ACCGCGTTAA
CTGGGAGTTG	CCATATCTGT	CAACCTGGGA	AACCGGCAAG	ACGCGGAAGC
CCGGCAGCAG	AAGCTGAGTT	TGTGCCATCA	TOCTGGAAGA	TGACCGCTCT
TCATGGAGTT	CAGCGAGCAC	CAACAACATC	AACCACAACA	CAGAGCTGCT
GACATCAGCT	CTGGACACCC	TGGTGGGGAA	AGGTCGCTTT	GCTGAGGTCT
GCCCATTGAG	GCTGAAGCAG	AACACTTCAG	AGCAGTTTGA	GACAGTGGCA
ATAAGGCCAA GTCAAGATCT	TTCCCTATGA	GGAGTATGCC	TCTTGGAAGA	CAGAGAAGGA
	GACATCAATC	TGAAGCATGA	GAACATACTC	CAGTTCCTGA
CATCTTCTCA	GCGGAAGACG	CAGTTCGGGA	AACAATACTG	GCTGATCACC
CGGCTGAGGA	CCAAGGGCAA	CCTACAGGAG	TACCTGACGC	GGCATGTCAT
GCCTTCCACG	GACCTGCGCA	AGCTGGGCAG	CTCCCTCGCC	CCGCGGATTG
CAGCTGGGAG	CAGTGATCAC	ACTCCATGTG	GGAGGCCCAA	GATGCCCATC
CTCACCTCCA	ACCTCAAGAG	CTCCAATATC	CTCGTGAAGA	ACGACCTAAC
GTGCACAGGG		GGCTTTCCCT	GCGTCTGGAC	CCTACTCTGT
CICCICCIC	TGTGACTTTG	AGTGGGCAGG	TGGGAACTGC	AAGATACATG
CTGTGGATGA	CCTGGCTAAC	CAGGATGAAT	TTGGAGAATG	CTGAGTCCTT
GCTCCAGAAG	TCCTAGAATC	CAUGATGAAT	CGTGCTCTGG	GAAATGACAT
CAAGCAGACC	GATGTCTACT	GAAGTAAAAG	ATTATGAGCC	TCCATTTGGT
CTCGCTGTAA	TGCAGTGGGA	CTGTGTCGAA	AGCATGAAGG	ACAACGTGTT
TCCAAGGTGC	GGGAGCACCC	ARATTCCCAG	CTTCTGGCTC	AACCACCAGG
GAGAGATCGA	GGGCGACCAG	AGGTTGACTG	AGTGCTGGGA	CCACGACCCA
GCATCCAGAT	GGTGTGTGAG		GANCGCTTCA	GTGAGCTGGA
GAGGCCCGTC	TCACAGCCCA	GTGTGTGGCA		AAGATTCCTG
GCATCTGGAC	AGGCTCTCGG	GGAGGAGCTG	CTCGGAGGAG	CAGGCTGGGC
AAGACGGCTC	CCTAAACACT	ACCAAATAGC	TCTTATGGGG	CHOOCIAGGC
ATGTCCAAAG	AGGCTGCCCC	TCTCACCAAA		

## FIGURE 3

MGRGLLRGLW	PLHIVLWTRI	ASTIPPHVQK	SVNNDHIVTD	nngavkppql
CKFCDVRFST	CDNOKSCHSN	CSITSICERP	QEVCVAVWRK	NDENITLETV.
CHDPKLPYHD	PILEDAASPK	CINKEKKKPG	etpphcscss	DECNDNIIFS
BEYNTSNPDL	LLVIPQVTGI	SLLPPLGVAI	SVIIIFYCYR	VNRQQKLSST
WETGKTRKLM	EFSEHCAIIL	<b>EDDRSDISST</b>	CANNINHNTE	LLPIELDTLV
GKGRFAEVYK	AKLKONTSEQ	PETVAVKIPP	YEBYASWKTE	KDIFSDINLK
HENILOPLTA	EERKTELGKQ	YWLITAPHAK	GNLQEYLTRH	VISWEDLRKL
GSSLARGIAH	LHSDHTPCGR	PKHPIVHRDL	KSSNILVKND	LTCCLCDFGL
SLRLDPTLSV	DDLANSGQVG	TARYKAPEVL	<b>ESRMNLENAE</b>	SPKQTDVYSM
ALVLWENTSR	CNAVGEVKDY	<b>EPPFGSKVRE</b>	HPCVESHKDN	VLRDRGRPBI
PSFWLNHQGI	<b>QHVCETLTEC</b>	WDHDPEARLT	aqcvaerfse	Lehldrlsgr

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# INTERNATIONAL SEARCH REPORT

Interpretated Application No.

PCT/US 92/09326

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